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(71) Applicants (for all designated States except US): MENTITO Y [FIFE]: FIN-12000 Parga FIN-12000 PIGMENTS OY [FIFE]: FIN-28840 Bjömeborg (f. 72) Inventors; and FIFE (f. 1200): (72) Inventors; and FIFE (for US only): ERIKSSON, [FIFE]: Managasan 16, FIR-10000 Karis (FI). Michael (FIFE): Legislants (f. FIR-1000): Karis (FI). Michael (f. 1200): FIR-28400 Urbis (f. 74) Agent: OY JALO ANT-WUORINEN AR; Stora Rob-4-6 A, FIN-00120 Helsingfors (FI).	Bo-Err ATTIL I). ertsgats	A ik t.
(54) Title: METHOD FOR TREATING FERROUS SULP	HATE	

(57) Abstract

The invention concerns a method for creating ferrous sulphate for the preparation of ferrous sulphate hydrate usable for chromium motion in cement manufacture. According to the invention, sulphuric acid containing ferrous sulphate obtained as a sic Optionate form titanium dioxide production and which contains at the most appr. 30 % by weight of sulphuric acid, in sentingliate with a Co-containing material, preferably cement, in an amount sufficient to give a pH-value to the end product which is 1.5 to 5, whereby the reaction temperature is allowed to increase to at the most 120 °C.

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WO 96/33133 PCT/F196/00198

Method for treating ferrous sulphate

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The object of the present invention is a method for treating ferrous sulphate for the preparation of ferrous sulphate hydrate which can be used in cement manufacture for the reduction of chromium.

As is known, cement is manufactured from limestone, silicate, iron and aluminium minerals together with optional fluorine and sulphur correction materials, which are ground and mixed homogenously and thereafter burned to form so called cement clinker. Cement also contains chromium compounds, which on contact with water, for example in concrete production, is converted to the Cr⁶⁺-form. This soluble chromate is undesirable as it can cause allergies in persons who come into contact with the cementwater mixture. It is known (e.g. EP 0054314), in order to reduce the Cr⁶⁺-content, to add ferrous sulphate to the dry cement clinker to reduce Cr⁶⁺ to Cr³⁺, which reduced

form is practically insoluble in the cement-water mixture. The added amount ferrous sulphate is generally in the range of 0.5 to 1 % by weight of the cement clinker.

Ferrous sulphate has a tendency to absorb water wherefore it is difficult to handle as such, for example to pump or 25 to blow. Ferrous sulphate is, however, commercially available e.g. in the form of its heptahydrate provided with an oxidation preventing coating, which coating imparts to the product also desirable flow characteristics. The use of such a coated preparation is, however, expensive ta-30 king into account the large amounts needed in cement manufacture. It has also been suggested (FI 842873) to use technical or commercial ferrous sulphate heptahydrate without said coating, whereby the ferrous sulphate heptahydrate first is subjected to drying by heating or ab-35 sorption using physical or chemical methods. Mixing of gypsum or fly ash in the ferrous sulphate is mentioned as a particularly suitable chemical method, whereas calcium

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oxide or cement alone are not said to give optimal results.

The problem underlying the present invention was to find a material for chromium reduction which, on the one hand, contains only such components that are "familiar" in cement manufacture and, on the other hand, are available at low cost and in large quantities, and the use of which in addition satisfies the environmental requirements one today places on the industry and its waste products. The problem according to the invention was solved by using, as chromium reduction agent, ferrous sulphate originating from the titanium pigment manufacture. The ferrous content in such a ferrous sulphate side product, which primarily in its monohydrate form, lies typically in the range of from 13 to 18 % by weight and it has a high sulphuric acid content, up to 30 % by weight, typically from about 10 to 30 % by weight, and is therefore extremely corroding and useless as such. This product contains in addition a considerable amount of moisture, up to 10 % by weight, and smaller amounts of trace elements. It is a product which is extremely difficult to handle and a burden on the environment, and has to be neutralized in order to be deposited. Such methods according to which the neutralization takes place in an aqueous solution to an alkaline pH, have been described, for example, in DE OS 4103311 and 3724677.

According to the invention, this moist and sulphuric acid rich ferrous sulphate obtained as a side product is neutralized and dried with a CaO-containing material, preferably cement, optionally together with limestone, to a pH-value of 1.5 to 5, thereby allowing the reaction temperature to rise to at the most 120 °C. In the reaction, a large part of both the free water and the water bound 35 in crystal water form leaves, and as major products ferrous sulphate hydrate, which is water soluble, primarily

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in the form of monohydrate, and gypsum, calcium sulphate, which is one of the starting materials in cement production, are obtained.

5 It is essential that the temperature does not exceed the said 120 °C in order to ensure that not all hydrate water leaves the product and that the iron is not oxidized to trivalent form which is useless for the chromium reduction. The product obtained has excellent handling, especially flow characteristics, and it can be pumped and blown in silos, and it does not dust, which is of essential importance in industry in outdoor handling.

Thus, according to the invention, a method has been provided according to which a waste product from the titanium dioxide pigment industry in a simple and economical manner kan be converted to a product which is easy to handle and which without any problems can be further utilized in full in cement manufacture. The product contains only such components which are already included among the cement manufacture materials and thus does not contribute with any side products whatsoever which are detrimental to the cement manufacture.

As CaO-containing material, cement or a cement containing 25 material, e.g. slag or fly ash cement, optionally together with limestone, are used. The CaO-containing material is added to the starting ferrous sulphate in substantially dry form. It is in principle possible to use any type of CaO-containing material, for example steel 30 slag, blast furnace slag, precipitator dust etc. provided they exhibit sufficient hydraulic characteristics. Products having a lower reactivity, e.g. filter dust from the cement and lime industry, limestone, and even fly ash, are usable in an initial stage of the process, but 35 in such case a post-treatment with more reactive material, e.g. cement, has to be carried out. It is also conWO 96/33133 PCT/F196/00198

ceivable to use hydrated and unhydrated lime, but the use of these extremely reactive materials require a strict temperature control.

- The amount of CaO-containing material to be added depends naturally on the sulphuric acid content in the starting material, but an amount of from 5 to 20 % by weight, calculated from the starting ferrous sulphate, is in most cases sufficient to neutralize to the desired pH and to 10 dry the product. The starting ferrous sulphate is used in the form it is obtained from the pigment production, that is in crystallized, filter dry pressed form, which generally means a moisture content of less than appr. 10 % by weight in the starting product. It is also possible to 15 carry out a careful pre-drying of the starting material. The addition of CaO-containing material leads to an exothermic reaction, which aids in the removal of free moisture from the material. Care has to be taken, however, that the temperature does not rise too much, for the abo-20 ve reasons, which conveniently can take place by regulating the amount and rate of addition of neutralizing agent, but also by suitably chosing the degree of fineness of the material.
- 25 The flow characteristics of the obtained product can be further improved by treating the product with fly ash, either already together with the neutralizing material or after neutralization, using an amount up to 20 % by weight of the ferrous sulphate. Fly ash as such does not possess sufficient hydraulic properties to function alone as both a neutralization and drying medium, but can very well be used as a post-treatment agent, or filler.

According to an advantageous embodiment of the invention, the neutralizing agent, preferably cement, optionally together with limestone, is added in an amount which gives a pH of appr. 3 to 4. It is also preferable to keep

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the temperature below appr. 105 °C, optimally at appr. 80 to 105 °C, whereby a soluble product suitable for passivation, is obtained. According to a preferred embodiment, the treatment can take place so that in a first stage limestone is added, which, as mentioned above, gives a slower reaction and a lesser temperature increase, for example to a pH of appr. 1.4, and thereafter the more active material, e.g. cement, is added to raise the pH to the desired pH end value. This procedure provides an excellent temperature control. If such a pre-treatment is carried out, the amount of cement or a material with comparative reactivity can be reduced, for example to 2 to 15 % by weight of the starting material. The pH value is measured by dissolving one part by weight of ferrous sulphate in 9 parts by weight of water.

The composition of the product obtained after the treatment varies to some degree depending on the composition of the starting material and the amount and type of neutralizing agents used. From the point of view of use, na-20 turally the ferrous content, i.e. the Fe2+-content. is of importance, and this is generally within the range of 10 to 18 % by weight, which corresponds to a ferrous sulphate content in pure hydrate form of appr. 30 to 55 % by 25 weight. In addition, the product typically contains appr. 15 to 30 % by weight of gypsum, CaSO, and moisture, typically 2 to 5 % by weight, and other products, e.g. cement and limestone. The product is well preserved during cement manufacture, i.e. there is no premature oxidation to Fe3+ in the mill, the product does not irritate and it is 30 compatible with the cement manufacture. The gypsum in the product can substitute part of the amount of gypsum normally used in cement manufacture.

35 The following examples illustrate the invention, without restricting the same. WO 96/33133 PCT/F196/00198

EXAMPLE 1

A test batch of appr. 1000 tons of raw ferrous sulphate hydrate from titanium dioxide pigment production which contained appr. 14 % by weight of Fe²⁺ and appr. 24 % by weight of sulphuric acid, was neutralized with cement (Partek Cement AB, Rapid) using an amount of 8 % by weight calculated from the ferrous sulphate hydrate. The mixing took place in a continuous concrete mixing plant. The mixing time was short, only a few seconds, whereafter the mixture was deposited for end reaction. Mixing took place without any disturbances from the materials used. An analysis of the end material showed that the Fe²⁺-content was substantially the same and that essentially no oxidation to Fe³⁺ had taken place. The pH of the end product was 1.5.

EXAMPLE 2

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Further test series have been carried out wherein raw 20 ferrous sulphate has been neutralized with different materials and also using different combinations. In the series 1 - 10, the ferrous sulphate according to the Example 1 was treated first with 4 and 8 % by weight of 25 limestone, respectively, and therafter with 6 and 2 % by weight of Rapid cement, respectively, calculated from the ferrous sulphate. In the series 11 to 22, raw ferrous sulphate was mixed with various types of CaO-containing material in an amount of either 10 or 20 % by weight, calculated from the ferrous sulphate. The pH was measured 30 (1 part by weight powder and 9 parts by weight water) after 1 day. The heat generation in the powder mixture was measured after a mixing time of 5 minutes. The results are given in the Tables 1 and 2.

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TABLE I		

5	Types	of limestone		B: Parfi C: Parfi D: Bypas		k Cement AB -
••	No.	Type of lime stone	ŧ	pН	Cement . %	рн
10			4	1.51	6	1.80
	1 2	A A	8	1.58	2	
	2	A	В	1.58	2	1.70
	3	В	4	1.58	6	1.81
15	4	В	8	1.64	2	1.72
		с	4	1.62	6	1.77
	5 6	č	8	1.67	2	1.73
		Ç	۰	1.0/	2	1.73
20	7	D	4	1.67	6	1.93
	8	D	8	1.73	2	1.86
	9	E	4	1.65	6	1.90

The results show that limestone in the amounts used, was not reactive enough to sufficiently raise the pH, but that a sufficient pH raise is obtained with a further addition of cement. When mixing the limestone, a very small heat generation was obtained for the samples 1 to 6, for the samples 7 and 9 the maximum temperature was 32 °C and for the samples 8 and 10, it was 40 °C. When mixing the cement, the maximum temperature for the samples 1, 3, 5, 7 and 9 was appr. 60 to 80 °C, and for the remaining samples appr. 40 °C.

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1.84

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TABLE 2

	No.	Additive	ŧ	Temp °C	. pH
5					
	11	Rapid-cement	20	-80	3.3
	12	Salmisaari			
		desulphuration product	20	~ 5	1.9
	13	Copper-slag sand	20	-30	1.8
10	14	Special steel slag (602 m2/kg)	ō	~50	1.7
	15	Ground steel slag (511 m2/kg)	20	~60	2.3
	16	Steel slag (unground)	20	~35	1.7
	17	Blast furnace slag (404 m2/kg)	10	69	
	18	Mason's cement	10	74	
15	19	Standard cement	10	78	2.1(3d)
	20	Quick cement	10	93	
	21	Standard cement +	10		
		fly ash	5	79	2.3(3d)
	22	Standard cement	15	81	2.7(3d)
20			-		

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In the samples 19, 21 and 22, the ferrous sulphate had been pre-treated with 8 % by weight of limestone. From the results it can be seen i.a. that the temperature can be regulated by means of a suitable choice of cement type.

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In the following Table 3. the pH and temperature increase (pH after one day and the maximum temperature of the mixture, respectively) in tests carried out by adding 10 % by weight of standard cement to three differently treated ferrous sulphate products, namely

1: ferrous sulphate treated with 8 % by weight of limestone

2: ferrous sulphate treated with 8 % by weight of Rapid cement

3: ferrous sulphate treated with 8 % by weight of limestone and thereafter with 8 % by weight of stan dard cement

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	No.	Max. temp.	Start.prod. pH	End prod. pH
	1	90	1.58	1.98
5	2	33	1.86	2.80
	3	22	2.95	4.04

The results show that for the products 2 and 3, a small or negligent temperature increase is obtained by means of the further addition of 10 % by weight cement, but that on the other hand the product 1 contains more free sulphuric acid.

EXAMPLE_3

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In this example, the capacity of the ferrous sulphate hydrate products made according to the process, namely products 2 and 3 from the Table 3 was measured. One part by weight of products 2 and 3, respectively, was mixed with 92 parts by weight of Rapid cement and 7 parts by weight of gypsum. The content of sixvalent chromium (Cr.6+) was measured in the obtained product. Also the reduction capacity of the product was measured using the standard SFS 5183 (24.3.86) by adding to the mixture an excess of 50 mg/kg Cr.6+ and measuring the amount of reduced chromium. Without the addition of extra chromium no measurable chromium content is obtained. The higher the value, the batter reduction capacity of the product

Sample	Cr ⁶⁺	Reduction capacity
2	0	39 mg/kg
3	0	37 mg/kg
		2 0

35 By adding amounts between 0.5 to 1 % by weight of the product obtained according to the invention to various commercial cement types, reduction capacities up to 49 mg/kg could be measured using the above mentioned standard method. The ferrous content (Fe²⁺) in the treated ŧ

cement varied typically between 0.01 to 0.2 mg/kg. In the following table, a compilation of analysis results is presented from tests where ferrous sulphate hydrate has been added to cement with varying starting chromium content.

TABLE 4

10	Cr ⁶⁺ P mg/kg	ddition %	Red. capacity mg/kg	Fe ²⁺ in cement
	24	0.7	6	0.07
	14	0.5	4	n.a.
15	25 (Luja d	0.5 cement)	13	n.a.
	25 (Rapid)	0.5	17	n.a.

20 n.a. = not analyzed.

Claims

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- Method for treating ferrous sulphate for the preparation of ferrous sulphate hydrate usable for chromium reduction in cement manufacture, characterized in that sulphuric acid containing ferrous sulphate obtained as a side product from titanium dioxide production and which contains at the most appr. 30 % by weight of sulphuric acid, is neutralized with a CaO containing material in an amount sufficient to give a pH value to the end product which is 1.5 to 5, whereby the reaction temperature is allowed to increase to at the most 120 °C.
- 2. The method according to the claim 1, characterized in that the CaO-containing material is used in an amount which gives an end pH value of 2.3 to 4.
 - The method according to the claim 1 or 2, characterised in that a temperature of at the most 105 °C is maintained.
 - 4. The method according to any one of the claims 1 to 3, characterized in that the CaO-containing material is cement.
 - 5. The method according to the claim 4, characterized in that limestone is used together with cement.
- 6. The method according to any one of the claims 1 to 5, characterised in that the starting ferrous sulphate is first treated with limestone or a smaller amount of cement, to a pH-value over appr. 1.4, and thereafter with cement to the end pH-value.
- 7. The method according to any one of the claims 1 to 5, characterized in that the starting ferrous sulphate is first treated with cement and optionally with limestone,

and thereafter with fly ash to improve the flow characteristics of the end product.

 The method according to any one of the claims 1 to 7,
 characterized in that the starting ferrous sulphate is subjected to drying prior to the neutralization.

INTERNATIONAL SEARCH REPORT

International application No.

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A. CLASSIFICATION OF SUBJECT M.	TTER	
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Documentation searched other than minimum do SE, DK, FI, NO classes as above	cumentation to the extent that suc	th documents are included in the fields searched
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Category* Citation of document, with indic		
X GB 1581539 A (LAPORTE 17 December 1980	INDUSTRIES LIMITED) (17.12.80), claims i	
	 ELSKABET AALBORG BRIK), 24 May 1984 (1-8
claim 1, abstract		
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GB-A-	1581539	17/12/80	CA-A- DE-A,A- FR-A-	50698 292637 108495 274502 236703 304649	77 57 23 30	31/01/80 05/04/79 02/09/80 20/04/78 05/05/78 26/04/78
WO-A1-	8401942	24/05/84	EP-A.A-	226018 012526 950182	57	04/06/84 21/11/84 01/11/84